# (19) World Intellectual Property Organization

International Bureau





(43) International Publication Date 8 July 2004 (08.07.2004)

**PCT** 

(10) International Publication Number WO 2004/057100 A1

(51) International Patent Classification<sup>7</sup>: 5/00, D06Q 1/12, G02B 5/128

D06P 1/00,

.

(21) International Application Number:

PCT/EP2003/012948

(22) International Filing Date:

19 November 2003 (19.11.2003)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

02425785.9

19 December 2002 (19.12.2002) EI

(71) Applicant (for all designated States except US): INDUSTRIA BERGAMASCA RIFRANGENTI S.R.L. [IT/IT]; Via Pederzola 1/3, I-24040 Scanzorosciate (BG) (IT).

(72) Inventors; and

(75) Inventors/Applicants (for US only): PARISI, Agostino [IT/IT]; c/o Idustria Bergamasca Rifrangenti s.r.l., Via Pederzola 1/3, I-24040 Scanzorosciate (BG) (IT). BARTOLI, Cesare [IT/IT]; c/o Industria bergamasca rifrangenti s.r.l., Via Pederzola 1/3, I-24040 Scanzorosciate (BG) (IT).

(74) Agent: GARAVELLI, Paolo; A.Bre.Mar. S.r.l., Via Servais, 27, I-10146 Torino (IT).

(81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PII, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

(84) Designated States (regional): ARIPO patent (BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

#### Declaration under Rule 4.17:

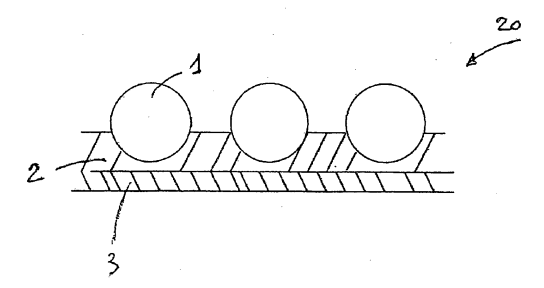
— of inventorship (Rule 4.17(iv)) for US only

#### Published:

with international search report

[Continued on next page]

(54) Title: PROCESS FOR MANUFACTURING RETROREFLECTIVE PRINTED MATERIAL



(57) Abstract: The present invention relates to a retroreflective printed product on fabrics a for use in clothing, sportswear, footwear and accessories with a lightreflecting effect, with good design flexibility which ensure trend-setting fashion sense as well as safety sense due to high visibility. More particularly, the invention deals with a continuous, flexible printing process without pollution and effluence, which use a printed mono- or multi-coloured design on paper base or other plastic base to be transferred over the surface of a layer (20) of glass microspheres (1) that are partially embedded in a support sheet (2) and coated with a very thin layer (5) of transparent thermo-adhesive bi-component polyurethane.

004/057100 A1

# WO 2004/057100 A1



 before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

### PROCESS FOR MANUFACTURING RETROREFLECTIVE PRINTED MATERIAL

#### DESCRIPTION

The present invention refers to a process for manufacturing retroreflective printed material.

It is known that retroreflective products used for safety garments can reduce risk of accidents, especially for some particular categories of people, such as, for example: firemen, paramedicals, adult and children playing sports.

The only commercial products suitable for reflective garments have generally been of the single-colored type. For example in US-A-4.763.98, US-A-5.283.101 and US-A-5.738.746 launderable retroreflective grey-colored products are disclosed. The following patents describe the possibility of obtaining colored effects and printed effects as well as reflective quality.

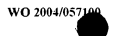
A retroreflective structure described in US-A-5.962.121 is capable of exhibiting a decorative effect both during the day and during the night, and particularly a rainbow-colored effect.

In US-A-4.605.461 a method is described for transferring a retroreflective pattern onto a fabric. Retroreflective images formed on garments and other substrates are described in US-A-4.102.562, while US-A-5.508.105 discloses a thermal printing system and a colorant/binder for printing frangible, retroreflective sheeting material. US-A-5.620.613 discloses printing of designs or emblems on garments, comprising a monolayer of

with a silk-screening system. When the prints of the first color are all dried, the subsequent colors can be printed through the same technique until the design on the layer of microspheres is completed. A similar patent for decorating textile surfaces, US-A-5.679.198, discloses a multi-step printing of many colors prepared with a polyester resin and an isocyanate hardener, dried before printing the following color and so on. Also in US-A-5.785.790 the same silk-screening multi-color printing technique is used with a system of colors made of polyester resin hardened with isiocyanate.

Many other patents (US-A-3.689.346, US-A-5.643.400, US-A-4.082.426, US-A-2.231.139, US-A-2.422.256, US-A-4.656.072, US-A-4.952.023) describe processes for producing retroreflective materials. US-A-6.120.636 discloses a high speed, low cost process for producing sheets patterned with drawings and emblems using a rotary screen printing system with cylinders and hardening with U.V. lamps.

Despite the above-described prior art situation, there still remain restrictive limits for printing retroreflecting products using many colors, with a high production speed, production flexibility and without ecological problems. From what is known, no one has previously found a practical way to produce a printed retroreflective product for fashion use using designs containing one or many colors. Some have proposed silk-screen printing with one water-based color or solvent-based colors but the above inventions are unfeasible for reproducing fashion designs with many colors upon a retroreflective material.



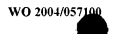
In the present invention as pointed out in Claim 1, a temporary support sheet is provided, with a monolayer of transparent glass microspheres partially embedded in a layer of softened polymer to a depth ranging between one-quarter and one-half of the microsphere diameter, as conventionally used in retroreflective materials, as described in US-A-3.700.305 and US-A-6.416.188. Then, after coating the layer of microspheres with a thin thermo-adhesive polymer film, a design is thermotransferred onto the microsphere surface.

Two kinds of commercial transfer-printed design may be used:

- (a) designs with sublimate pigments printed on a paper base; or
- (b) designs having a polymer film supported by a release paper base or a polymer film base, such as for example a film of polypropylene.

In case of transferring a printing with sublimate pigments (a), the transfer temperature ranges between 180°C and 220°C. A temperature close to 220°C causes a maximum yield of color transferring, but also a partial transferring of colors at lower temperatures may give a satisfactory aesthetic design on the final retroreflective product.

In case of transferring a printed polymer film as shown in (b), the layer of microspheres is beforehand coated with a thin layer of bicomponent polyurethane. The thin layer of polyurethane resin dried but not cured operates as thermo-adhesive between microspheres and printed film. In this case, the print transfer temperature is lower than 150°C, and preferably between 100°C and 120°C.



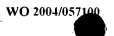
As regards the above-described prior art, many patents use the screen-printing technology (US-A-5.620.630, US-A-5.785.790 and others). With this printing system, it is concretely impossible to print designs containing many colors while maintaining the design accuracy and the perfect fitting of various printed colors, not as is normally done on a textile support but on a layer of microspheres to produce retroreflecting materials. The same considerations can be done with a rotary screen-printing system (US-A-6.120.636).

The present invention instead provides a flexible, ecological, easy-to-apply process, for obtaining printed retroreflective products especially, but not restrictively, for fashion garments where rich designs and colors are demanded and appreciated. The printing transfer machine needs a low-cost investment compared with other printing processes; no auxiliary equipment and small floor space are required, and no pollution or obnoxious effluence is produced. Moreover, the availability of commercial transfer printed papers is considerable.

A special feature of the present invention is the possibility of vacuum application of a thin aluminium reflecting layer after the printing process. In this case, it is possible to avoid the application of a transparent dielectric mirror though maintaining a sufficient reflective intensity for a printed fashion product.

Over the printing or over the reflective aluminium layer, a polyurethane two-components resin is coated, dried and laminated over a fabric. The polyurethane resin coating may be substituted with a thin layer of a hot-melt adhesive being applied.

The present invention will be better described by some preferred embodiments thereof, provided as a non-



limiting example, with reference to the enclosed drawings, in which:

- Figure 1 shows a schematic sectional view of an article of clothing 10 at the final stage of production according to the present invention;
- Figure 2 shows a schematic view of a continuous apparatus for doctor blade on roll coating of a supported layer of microspheres;
- Figure 3 shows a schematic view of a production machine for transferring printed designs using sublimation pigments;
- Figure 4 shows a schematic carrier web, which secures microspheres thereon in a desired temporary arrangement;
- Figure 5 is a plan view showing a schematic design of a printed paper; and
- Figure 6 schematically shows the drawing transfer of Fig. 5 from the original printed sheet to the surface of the layer of microspheres.

In the invention as described and shown, a specific terminology is used for better clarity. However, the invention is not constrained to the specific terms being chosen and it is obvious that every chosen term comprises every technical equivalent that generates a similar behaviour.

Fig. 4 is a cross sectional view of a carrier web 20, which secures glass microspheres 1 on a temporary transport support. The carrier web used as a sheet material is produced as described in US-A-4.102.562. The microspheres 1 used in the present invention typically have an average diameter in the range of about 30 to 200 microns and a refractive index of about 1.7 to 2.0. Preferably the glass microspheres 1 are arranged



substantially in a monolayer on a temporary carrier sheet 20, which comprises a backing sheet 3 and a polymeric coating film 2. The polymeric coating 2 is a softenable material such as polyethylene, polypropylene and the like. The stiff backing sheet 3 could be kraft paper, polyester film and the like.

The microspheres 1 may be arranged upon the temporary carrier sheet 20 by printing, cascading, transferring, and screening or any convenient transfer process.

The microspheres 1 can be embedded in the carrier sheet 20 with a pressure roll or by heating the softened polymer, to a depth between about 20% to 40% of their average diameter.

Fig. 1 shows a sectional view, not to scale, of a portion of an article of clothing 10 that is partially delaminated from the carrier web comprising the polymeric coating 2 and the kraft paper or polyester film backing 3.

Disposed adjacent to the non-embedded glass surface of the microspheres 1 is a transparent dielectric mirror 4, a priming layer 5 of bi-component polyurethane of about 1 micron.

The layer 6 reduces the printed layer, whose thickness is less than 0.1 microns, in the case of sublimate pigments (a) and less than 0.5 microns in the case of transfer printing supported by a polymer film (b).

With reference again to Fig. 1, the printed design over the microspheres 1 is covered with a layer 7 made of vacuum-nebulised aluminium, or other light reflecting material. Obviously, in this case the layer of transparent dielectric mirror 4 is not necessary.

With reference again to Fig. 1, finally, a binder layer 8 will provide an adequate thermal adhesion with a base fabric 9, for example a polyester/cotton fabric, a



nylon knitted fabric made of a Lycra® or other textile bases.

Fig. 2 and 3 are schematic drawings of apparatus used in the invention, which include a well-known rotary machine 29 for thermal transfer printing of the calender type (manufactured by Lemaire, Roubaix, France or Monti Officine, Thiene, Italy).

The composite microspheres layer 33 (supplied by cylinder 40), as described in Fig. 4, together with the printed paper 30 (supplied by cylinder 24) are pressed between heated cylinder 27 and felt 26 in a continuous process (Fig. 3). Out of the machine, the paper 31 without the design is wound on cylinder 25 on one side, and the printed layer of microspheres 34 is wound on cylinder 32 on the other side.

In Fig. 2 the continuous printing process is made on the composite material 33 (supplied by cylinder 40) coated (in machine 23) with a polyurethane layer 5 (supplied by cylinder 22) as shown in Fig. 3. At the end of the process, a product 34 is obtained that is wound on cylinder 28.

Fig. 5 is a schematic plan view showing a transfer paper 30 printed with nature image containing 8 colors a, b, c, d, e, and f. The commercial offer of transfer printed paper is remarkable. This type of paper is widely used in many applications in textile industries but also in several areas such as accessories, furniture, interior decorations, motor vehicles and the like.

Samples of the present invention have been prepared using transfer printed papers from Transfertex GmbH, Kleinostheim, Germany and a special polypropylene printed film Decotrans® from Miroglio Sublitex, Alba, Italy.

Fig. 6 is view of partially removed released paper 31 without the design from the carrier web, which secures microspheres covered with the printed transferred image 34.

The invention will be further explained by the following illustrative examples, which serve the purpose of showing the features and advantages of this invention. However, the ingredients and the specific amounts recited therein, as well as other conditions and details are intended to be not limiting of the scope of the present invention. Unless otherwise specified, all amounts are expressed in the examples are in parts by weight.

#### EXAMPLE 1

Cascading the microspheres on a Kraft paper covered with an acrylic auto-adhesive film produced the monolayer of glass microspheres having diameters between 40 and 100 microns. The layer of microspheres was then transferred onto a support sheet of polyester covered with a low-density polyethylene thermo-adhesive film of 50-micron thickness. The transfer was made with a heated calender as shown in Fig. 3 at a cylinder temperature of 140°C. The contact time was 5 seconds and the pressure between the cylinder and the felt was 5 bars, in order to obtain a penetration of the microspheres onto the polyethylene film of about 40% of their diameter. The exposed surface of the microspheres was then coated with a transparent dielectric mirror as described in US-A-3.700.305.

A bi-component polyurethane priming layer was next applied over the electric mirror, by coating a solution of the following formulation 1 with a doctor knife-coating machine or a graved-roll coating machine:



Ingredients	Parts	by
	Weight	
Polyurethane resin	100	
("B 10" from Coim)		
Curing agent	5	
("Imprafix TH" from		
Bayer)		
Methylethylketone	150	
Formulation 1		

The resin has been dried and partially cured at 110°C. The amount of transparent film layer is about 4  $g/m^2$ .

At the end of the oven as described in the Fig. 2, the product is running into the calendar heated at 130°C and laminated with the printed polypropylene Decotrans® design shown in Fig. 5. The contact time is about 10 sec. Then the polypropylene without the design and the printed microspheres were separately unwound.

Subsequently, a solution of the following polyurethane formulation 2, using a knife on roll coating, was coated over the printed layer at approximately a 125-micron thick wet substance:



Ingredients	Parts	рy
	Weight	
Polyurethane resin	100	
("B 10" from Coim)		
Curing agent	5 ·	
("Desmodur RFE"	•	
from Bayer)		
Methylethylketone -	40	
Melamine curing	3	
agent ("C6" from		
Coim)		
Formulation 2		

The resin has been dried at 80°C. At the end of the oven the surface of the resin was superposed and calendered on a white polyester/cotton fabric containing 65% of polyester and 35% of cotton. After calendering the laminated compound at 100°C and a pressure of 5 bars, the compound was cooled and the polyester film was peeled off. Subsequently the printed retroreflective textile was cured at 150°C for 2 min.

#### EXAMPLE 2

A monolayer of glass microspheres having similar characteristics as those mentioned in Example 1 was deposited onto the low density 50-micron polyethylene film supported by a 40-micron polyester carrier. The glass spheres-covered carrier was then heated for 2-4 min at 150°-160°C and penetrated into the softened polyethylene. The glass microspheres thus became embedded in polyethylene for about 40% of the sphere diameter and formed a monolayer therein with little or no space between spheres. The coating with a transparent dielectric mirror



and the subsequent steps of production were the same as described in Example 1.

#### EXAMPLE 3

The monolayer of glass microspheres having diameters between 40 and 100 microns was produced by cascading the microspheres onto a thick release paper covered with an acrylic auto-adhesive film as described in Example 2 of US-A-4.075.049. The resulting microspheres binder composite was doctor-knife coated with a water polyether polyurethane dispersion having the following formulation 3:

Ingredients	Parts	by
	Weight	
Polyurethane water	100	
based resin		
("Idrocap 930" from		
Icap)		
Curing agent	5 .	
("Icaplink X3" from		
Icap)		
Water	40	
Thickening agent	a.r.	
("Idrocap 200" from		
Icap)		
Formulation 3		

The amount of wet resin was about 10  $g/m^2$  and was adjusted with the doctor knife profile, resin dilution and viscosity. The amount of dry film was about 3  $g/m^2$ . The resin was partially cured at 110°C.

At the end of the oven as described in Fig. 2, the product was run into the calender heated at 130 °C and



laminated with the printed polypropylene Decotrans® design shown in Fig. 5. The contact time was about 10 sec. Then the polypropylene without the design and the printed microspheres were unwound. The resulting printed composite was worked according to whether it comes covered or not covered with a vapour coating of a metal such as aluminium light reflecting material 7 in Fig. 1.

In case the composite was metallised, the subsequent process was the same as described in Example 1. In case the composite was not metalleisd, the subsequent treatment was polyurethane knife coating and textile lamination.

printing effect without reflecting light aluminium is very regular but the average reflectivity was between 8 and 15 cd/(luxm), that was a low value for a technical product but that remained effective for a fashion fabric. The metal layer of the printed retroreflective fabric that was metallised favourably affects the design colors and reflectivity is greater than 50 cd/(luxm).

#### EXAMPLE 4

The monolayer of glass microspheres having diameters between 40 and 100 microns was produced by cascading the microspheres onto a thick release paper covered with an acrylic auto-adhesive film as described in Example 2. The exposed surface of the microspheres was then coated with a transparent dielectric mirror. Then, the transfer print process was made using a commercial transfer printed design with sublimate pigments (a) from Transfertex GmbH, Kleinostheim, Germany. The transfer temperature was about 185°C. In fact the heated roll was in contact with the back of the transfer paper, therefore the real temperature of the glass layer was higher than the real temperature of the printed paper but sufficient for obtaining a good

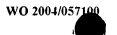


yield of pigments sublimation onto the upper surface of the microspheres. The composite material was metallised and coated using Formulation 2 with a knife on roll coating machine. The resin dried at 80°C. At the end of the oven the surface of the still tacky resin was superposed and calendered on a white polyester/cotton fabric containing 65% of polyester and 35% of cotton. After calendering the laminated compound at 100°C and a pressure of 5 bars, the compound was cooled and the polyester film was peeled off. Subsequently the printed retroreflective textile was cured at 150°C in an oven for about 2 min for finally curing the resin.



#### CLAIMS

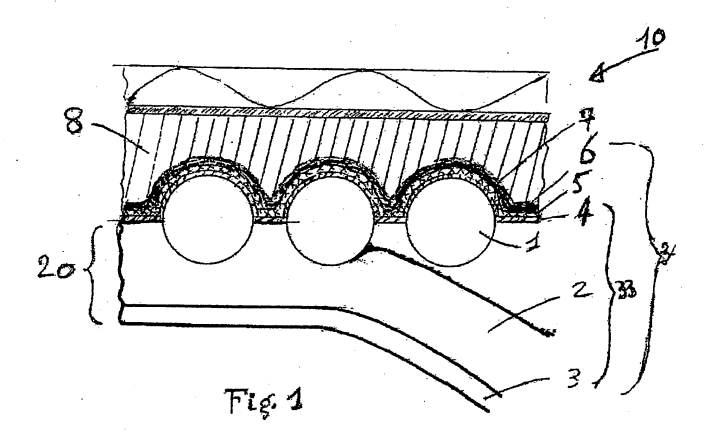
- 1. Process for manufacturing retroreflective printed material with a flexible printing process, without using pollution products and effluence, said process comprising the steps of:
- (a) partially embedding, in a support layer (2) of paper or polymer foil, a monolayer (20) of transparent glass microspheres and coating a reflecting layer (4) over a free surface of the microspheres (1);
- (b) applying a thin coating layer (5) of a dry bicomponent polyurethane product;
- (c) while the polyurethane layer (5) is partially cured, applying a transfer printed support (6) and transfer an image from the support (6) to the microspheres (1) with a calender (29);
- (d) laying an aluminium film (7) by vacuum deposition;
- (e) coating a bi-component polyurethane layer (8) of a wet substance and drying the layer (8);
- (f) while the polyurethane layer (8) is partially cured,
  applying a support fabric (9);
- (g) stripping away the support layer (2) of the microspheres (1) and curing the fabric (9) covered with the printed microspheres (1).
- 2. Process for manufacturing a printed retroreflective material according to claim 1, characterized in that the printed design to be transferred is a sublimate pigments design printed on a paper base.
- 3. Process for manufacturing a printed retroreflective material according to claim 1, characterized in that the printed design to be transferred is a printed design on a polymer carrier film supported by a release paper or a polymer base.

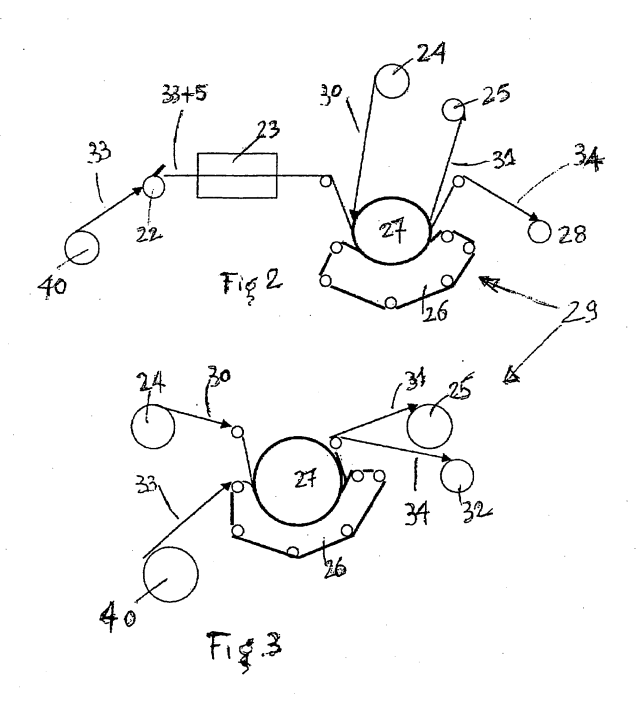


- 4. Process according to claim 1, characterized in that the layer (20) of microspheres (1) layer is deposited upon an acrylic auto-adhesive film (2).
- 5. Process according to claim 1, characterized in that the layer (20) of microspheres (1) layer is deposited upon a polyethylene thermo-adhesive film (2).
- 6. Process according to claim 1, characterized in that:
- (a) the first layer of polyurethane is a water-based dispersion polyurethane and the second polyurethane layer is a bi-component solvent solution polyurethane, in case an acrylic auto-adhesive film (2) is used for the microspheres (1),
- (b) the first and second layer of polyurethane are in a solution of organic solvents if a polyethylene thermo-adhesive film (2) is used for the microspheres (1).
- 7. Process according to claim 1, characterized in that the temporary layer (20) of microspheres (1) is directly thermo-printed with sublimate pigments designs without alternatively the step of coating with a dielectric mirror (4) or the step of vacuum deposition of the aluminium film (7).
- 8. Process according to claim 7, characterized in that, when the coating with aluminium (7) is omitted, the second polyurethane coating is colored so to obtain an aesthetic combination between the printed image and the colored polyurethane resin.
- 9. Process according to claim 1, characterized in that said microspheres (1) are embedded into said polymeric layer (2) down to a depth included between 40% and 50% of their average diameter.
- 10. Process according to claim 1, characterized in that said thin layer (5) is equal to about  $4 \text{ g/m}^2$ .

- 11. Process according to claim 1, characterized in that said step (c) of transferring the support (6) to the surface of the microspheres (1) with a calender (29) is carried out at a temperature included between 100°C and 180°C.
- 12. Process according to claim 1, characterized in that said bi-component polyurethane (8) is coated with a thickness of about 125 microns of a wet substance and is dried at about  $80^{\circ}$ C.
- 13. Process according to claim 1, characterized in that said step (g) of curing is carried out at a temperature of about 150°C.

# 10/540098





# 10/540098

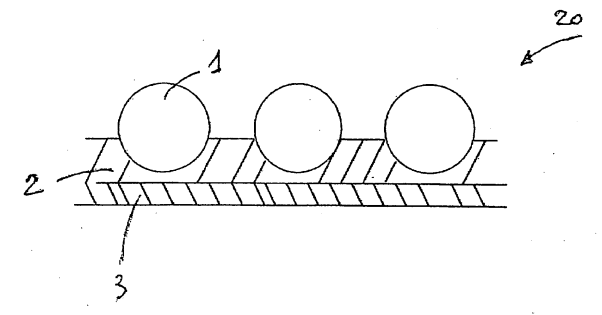
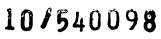
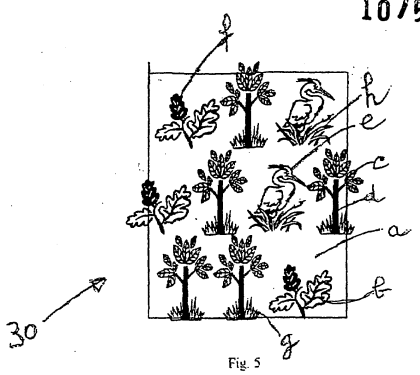
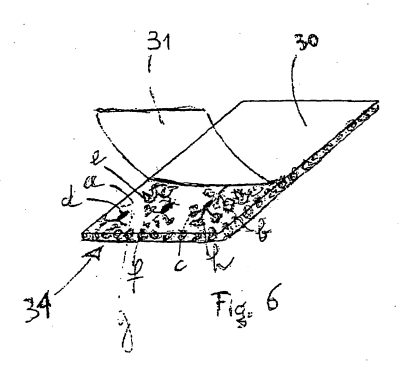


Fig. 4







# RNATIONAL SEARCH REPORT

onal Application No PCT/EP 03/12948

CLASSIFICATION O IPC 7 D06P1/00 D06P5/00 D0601/12 G02B5/128 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 7 DO6P D06Q G02B B41M A41D Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, WPI Data, PAJ C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category ° Relevant to claim No. Α US 4 153 412 A (BAILEY TERRY R) 1-13 8 May 1979 (1979-05-08) example 2 A WO 94/25666 A (MINNESOTA MINING & MFG) 1 - 1310 November 1994 (1994-11-10) cited in the application examples Α WO 92/07990 A (REFLEX PROMOTION 1-13 INTERNATIONAL) 14 May 1992 (1992-05-14) cited in the application example Α WO 96/24867 A (MINNESOTA MINING & MFG) 1 - 1315 August 1996 (1996-08-15) examples Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents: T later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not considered to be of particular relevance cited to understand the principle or theory underlying the "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an invention step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. \*O" document referring to an oral disclosure, use, exhibition or document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 26 April 2004 04/05/2004 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,

Fiocco, M

Fax: (+31-70) 340-3016

### INTERNATIONAL SEARCH REPORT

Information on patent family members

Intel Onal Application No
PCT/EP 03/12948

Patent document		Publication		Potent Corili	D 1.11
cited in search report		date		Patent family member(s)	Publication date
US 4153412	Α	08-05-1979	NONE		
WO 9425666	Α	10-11-1994	US	5344705 A	06-09-1994
			US	5503906 A	02-04-1996
			CA	2160361 A1	10-11-1994
			CN	1122621 A ,B	15-05-1996
			DE	69408978 D1	16-04-1998
			DE	69408978 T2	29-10-1998
			EP	0697042 A1	21-02-1996
	•		ES	2113657 T3	01-05-1998
			JP	8509783 T	15-10-1996
			WO -	9425666 A1	10-11-1994
			US	5916399 A	29-06-1999
			US	5620613 A	15-04-1997
WO 9207990	A	14-05-1992	DK	257390 A	26-04-1992
			US	5679198 A	21-10-1997
			AU	8851791 A	26-05-1992
			CA	2094708 A1	26-04-1992
			DE	69125174 D1	17-04-1997
			DE	69125174 T2	16-10-1997
			MO	9207990 A1	14-05-1992
			EP	0561793 A1	29-09-1993
			JP	3066979 B2	17-07-2000
			JP	65093 <b>9</b> 4 T	20-10-1994
•		•	KR.	228593 B1	01-11-1999
			US	5510178 A	23-04-1996
			US	5612119 A	18-03-1997
			US	5785790 A	28-07-1998
WO 9624867	Α	15-08-1996	US	5706133 A	06-01-1998
			AU	4909496 A	27-08-1996
			CA	2210493 A1	15-08-1996
			CN	1173923 A	18-02-1998
	•		DE	69628283 D1	26-06-2003
			DE	69628283 T2	01-04-2004
			EP	0808471 A1	26-11-1997
			JP	11500236 T	06-01-1999
-			WO	9624867 A1	15-08-1996